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ABSTRACT

Method and apparatus for delivering aerosolized medication employing a variable-volume device and a carrier nozzle.

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Details of Associated Provisional Applications Not:

The following statement is a full description of this invention including the best method of performing it known to us.

**METHOD AND APPARATUS FOR DELIVERING
AEROSOLIZED MEDICATION**

Field of the Invention

The present application, which is a divisional application derived from Application No. 992179, relates to methods and apparatus for delivering a dose of aerosolized medication for inhalation by a patient into the lungs.

Background of the Invention

Devices are increasingly being used for delivering medication for therapeutic treatment of the lungs. For example, in the treatment of asthma, inhalers are commonly used for delivering medications such as β_2 agonists and anti-inflammatory agents such as corticosteroids. Two types of inhalers are in common use, metered dose inhalers (MDIs) and dry powder inhalers (DPIs). Both types have their effects on delivery of medication, which is typically in the form of a small particle or powder, into the airways of the lungs at the location of the medication being treated.

In the MDI device, the medication is provided by the pharmaceutical manufacturer in a pressurized aerosol canister, with the medication being suspended

or dispersed in a liquid propellant such as a chlorofluorocarbons (CFC) or hydrofluorocarbons (HFC). The customer includes a dosing valve having a bottom discharge area which can be depressed forward from the customer to discharge a controlled volume of propellant-medicament mixture in the form of an aerosol. Inspecting the nozzle of propellants in which particles of the medicament are suspended or dispersed, a typical MDI or can will such a customer includes a dosing valve as shown and means. The customer is known how the dosing valve or bottom discharge area of the customer being mounted in a base of the nozzle. Depressing the base out of the customer causes the aerosol to be pushed forward from the customer to form a controlled volume of medicament in discharge through the nozzle.

The dosing valve defines a passage in fluid communication with the nozzle, the passage leading in series to a nozzle-like portion of the dosing, such that the medicament medicament may be released after it exits the nozzle-like portion. The patient either knows the nozzle-like portion has the base of the base around the nozzle-like, or holds the nozzle-like at a slight distance away from an open mouth. The patient thus depresses the nozzle to discharge the medicament, and disbursement starts.

Existing MDIs suffer from a number of significant disadvantages. One problem with existing MDIs is poor delivery efficiency of the medicament. It has been estimated that on average, with existing MDIs, only about 10 percent of the

medication is delivered to the mouth and throat. Ideally, the velocity of the aerosol should match the velocity of the patient's inspired breath so that the particles are entrained in the breath and carried into the lungs. With many existing MDIs, the exit velocity of the aerosol substantially exceeds the velocity of the patient's breath. The high-exit velocity places added on the back of the throat, causing irritation and sticking.

The author later concurring in the poor delivery efficiency of existing MDIs is excessive length of the plume or length of time exiting the device. In existing MDIs, this length typically exceeds 20 centimeters, which makes it difficult for the patient to inhale the exiting breath.

In an effort to decrease plume velocity, some MDI designs have added obstacles between the aerosol nozzle and the nozzle-like. Although spaces improve delivery efficiency, most of the drug vehicle is discharged from the nozzle before and exists on lower surfaces of the spaces, and is therefore susceptible to inhalation by the user. Thus, little vital passes and suffer from unacceptable low delivery efficiency.

Furthermore, although dry powder inhalers inherently avoid some of the aforementioned problems of MDIs, such as excessive aerosol velocity, DPDs suffer from the problem of impacting and sticking of medicament on the lower surfaces of the device, particularly under moist environmental conditions such as high relative humidity, which leads to coarse particle aggregation.

medication dose which is dispensed from the nozzle actually reaches the lungs where it can achieve the intended result.

Poor delivery efficiency is caused by a number of factors. One of these is incomplete dispersion of propellants, resulting in a large portion of the aerosol dose being delivered in a form which cannot be taken into the lungs. For effective delivery of medicinal medicament to the lungs of the lungs, it is believed that most of the particles within are required to be less than 10 microns (one micron-one-thousandth of a millimeter) in size, and preferably less than 1 micrometer and 5 microns. Insufficient dispersion of propellants in the nozzle of the nozzle results in a reduced fraction of the aerosol dose being delivered to the form of relatively large liquid droplets instead of the dry particle aerosol form. Such droplets cannot be inhaled, but rather tend to impact the walls of the nozzle and at the back of the patient's throat, with the result the much of the medicament is swallowed. The local concentration of medicament in the mouth and throat can cause local immune-suppressive responses, as well as development of drug tolerance in the case of corticosteroids. Additionally, protracted & frequent course of the aerosol nozzle of the pressurized can, which decreases acceptability and safety of the aerosol. Further, the aerosol medication has been estimated to cost U.S. patients about \$750 million per year.

Another factor contributing to the problem of poor delivery efficiency is high throat velocity of the aerosol as it exits the nozzle-like, which tends to lead to

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Another problem with existing MDIs is the difficulty patients have in coordinating their inhalation with the discharge of the aerosol. As currently operated MDIs, patients frequently inhale too early or too late to effectively inhale the medicament. Although a number of breath-activated MDIs have been devised to address this problem, most of these devices suffer discharge at the very onset of the patient's inspiration effort. Depending on the long duration being required and to breathe, it may often be more desirable for the medicament to be discharged near the peak of the patient's inhalation rather than the beginning. Further, it may be desirable to be able to selectively vary the point in the patient's inhalation at which medicament is discharged in order to refine the location of drug delivery in the respiratory system. These advantages are not possible with existing MDIs.

Accordingly, it has been an object of the present invention to provide a method and apparatus for delivering an aerosolized medicament in which the relative velocity of the aerosol or the exit of the apparatus approximately matches the velocity of the patient's inspiratory breath. Another object of the present invention is to provide a method and apparatus for delivering an aerosolized medicament in which the relative velocity of the aerosol or the exit of the apparatus approximately matches the velocity of the patient's inspiratory breath.

It has been another object of the invention to minimize dispersion and mixing of the drug particles in the base of an aerosol within an inhaler apparatus.

It has been a sole further object of the present invention to provide a method and apparatus for delivering an aerosolized medication in which the length of the tube of aerosolized medication which exits the apparatus is as short as possible.

A further object of the invention has been to provide a method and apparatus for maximizing the dispersion of liquid propellants in an inhaler.

Still another object of the invention has been to provide a method and apparatus for delivering an aerosolized medication in which impaction and sticking of medication on the inner walls of the apparatus is minimized.

It has been another object of the present invention to provide a method and apparatus for delivering an aerosolized medication in which the discharge of medication is synchronized with the patient's inspired breath, and in which the timing of the discharge is relative to the patient's breath can be independently varied.

Summary of the Invention

The above and other objects of the invention are achieved by the methods and apparatus of the invention in which three novel techniques and devices are used to prevent sticking of the propellant-medication mixture with air or increase dispersion of propellants to slow down the aerosol plume before it reaches the exit of the apparatus, and to reduce the impaction of aerosol on the inner walls of the apparatus. The invention also provides an apparatus and method for synchronizing the actuation of the cylinder with the patient's respiratory effort control on the dispensing of the apparatus.

In one embodiment of the invention, the apparatus is configured so that the nozzle discharge outlet directs a plume toward the open end of the nozzlepiece. The air tube is arranged to allow an air jet away from the open end of the nozzlepiece as to impinge on the plume. The air tube is supported, which is cradled by one or more hollow spacers connected to the wall of the nozzle, with the hollow passage of each spacer being connected at one end to a corresponding passage through the nozzle wall to enable air inside the nozzle and at the other end to the tube of the air tube. When the patient inhales on the open end of the nozzlepiece, air is drawn into the air tube to cause an air jet to exit the air tube. Once this air jet has been established, the cylinder is actuated to discharge a plume of aerosol toward the air jet.

The plume and air jet meet, causing mixing and dispersion of the plume.

In another embodiment of the invention, the nozzle is positioned to direct a plume away from the open end of the nozzlepiece toward the end of the nozzle, which end is automatically closed by an end cap. The air tube is inserted on the end cap, with the tube of the air tube connected to a passage through the end cap to enable air outside the nozzle. Utilizing by a patient on the open end causes air to be drawn through the air tube in a direction toward the patient's mouth. Once the air jet from the air tube has been established, the cylinder is actuated to direct a plume toward the closed end of the nozzle. The air jet and plume meet, causing mixing and dispersion of the plume. The plume exits directly toward the patient.

More specifically, the invention provides a aerosol dose inhaler apparatus including a housing adapted to support a pressurized cylinder, the housing having an actuator and nozzle assembly with a tube adapted to receive the nozzle ends from the cylinder, the housing further including a generally planar nozzle

- 3 having an open and closing a nozzlepiece adapted to be inserted into the nozzle of a tube, a nozzle discharge outlet of the actuator and nozzle assembly being positioned to direct a plume of aerosolized medication from the nozzle and an air tube supported within the nozzle and having an air tube inlet in fluid communication with ambient air outside the nozzle, the air tube being oriented so that air flowing out of the air tube outlet is directed so as to impinge on a plume of aerosolized medication discharged from the cylinder through the nozzle discharge outlet. Then, an inhaler effort control on the nozzlepiece causes air to flow into the air tube inlet and out the air tube outlet to impinge on the plume and thereby enhance dispersion and mixing of the medication within the nozzle. The air jet from the air tube then causes the plume to slow down so that the velocity of the aerosol exiting the device opportunity reduces the velocity of a patient's inspired breath. Slowing down the plume also increases the residence time of the aerosol within the apparatus and tends to a shorter time to be inhaled. The increased mixing and residence time prevent dose dumping.
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utilizing the nozzlepiece, so that the total length of nozzle is very short, thereby further increasing residence time of the aerosol within the device.

- To reduce impaction and sticking of medication on the inner walls of the apparatus, the invention provides an aerosol flow control apparatus, useful for other ACM or DNF devices, including a housing defining a nozzle, the nozzle having an open end defining a nozzlepiece and a substantially closed end defined by an end wall remote from the nozzlepiece, with a medication dispensing assembly being arranged within the housing to direct medication from the nozzle. The medication dispensing may be a pressurized cylinder with actuator and nozzle, or alternatively may be a dispenser for medication in dry powder form. The end wall includes a plurality of auxiliary air tubes in fluid communication with ambient air outside the nozzle, the auxiliary air tubes opening from the nozzle adjacent the inner wall of the nozzle. A direction generally toward the open end of the nozzlepiece. The nozzle further includes a plurality of venturi passages connected to the inner wall thereof.
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downstream of the nozzle air tubes, the auxiliary air tubes and venturi passages comprising to establish a venturi air flow along the inner wall of the nozzle open to inhaler effort being exerted on the nozzlepiece. The auxiliary air flow acts as a buffer or boundary layer flow along the inner wall of the nozzle, reducing the likelihood of aerosol droplets or dry powder impacting and permanently sticking to

which are angled at an angle to the initial direction in which the patient would inhale.

The breathing device provides an assisted flow control apparatus for use with a prescribed volume of medication, in which discharge of the aerosol plume is caused by the patient's inspiratory effort, with no closing of the discharge area in relation to the inhalation being selectively variable. To these ends, the apparatus includes a housing adapted to support the canister between a first position in which the discharge area of the canister is in an inspiratory position to a second position in which the discharge area is in an expiratory position for discharging a measured volume of medication, the breathing device including an outlet through which a user can intake, the outlet defining a primary air passage. A canister rotatable is arranged in the housing and is rotatable from a first position in which relative movement between the canister body and discharge area is prevented to a discharge position in which such movement is permitted. The canister rotatable has a port at, or alternatively is, attached to, a stalk such as a balloon or a variable discharge plume assembly which defines a variable-volume chamber. The stalk includes a resilient member which urges the canister into the second position upon movement of the canister rotatable into discharge position. A secondary air passage extends through the housing between the primary air passage and outside of said stalk or housing, the secondary air passage including a valve. The variable-volume chamber is in fluid communication with a plenum of the canister, whereby inhalation of a user through the

outlet causes a flow pressure in the variable chamber so as to evacuate air from the chamber and thereby cause the canister member to move into the discharge position.

By appropriate selection of design parameters such as the chamber cross-sectional area, the force exerted by the resilient member on the canister, the vessel size, and

- 3 On secondary air passage discharge, the device can be designed to cause inhalation of the canister near the path of a patient's inspiratory effort.

The device preferably further includes means for selectively varying the timing of inhalation. For instance, the device may include an adjustment device indicating how the secondary air passage is set as a variable flow restrictor. Tuning

- 10 the flow rate selectively increases the amount of flow available, such that for a given inspiratory rate through the mouthpiece, the amount of time required to evacuate the chamber sufficiently to cause inhalation is improved. Conversely, tuning the valve to the opposite direction decreases the amount of time required to cause inhalation.

These and other objects and advantages of the present invention shall

- 13 become apparent from the accompanying drawings and the description thereof.

Detailed Description of the Invention

The accompanying drawings, which are incorporated in and constitute a part of this specification, illustrate various embodiments of the invention and, together with the general description of the invention given above and the detailed description given below, serve to explain the principles of the invention.

FIG. 1 is a perspective view of an inhaler in accordance with the principles of the present invention.

FIG. 2 is an exploded view of the inhaler of FIG. 1.

FIG. 3 is a cross-sectional view of the inhaler taken along lines 3-3 of

FIG. 1.

FIG. 3A is a partial cross-sectional view showing an alternative embodiment of the canister and stalk of the inhaler.

FIG. 4 is a cross-sectional view similar to FIG. 3, showing an alternative embodiment of the inhaler.

FIG. 2.

FIG. 5 is a cross-sectional view similar to FIG. 3, showing yet another alternative embodiment of the inhaler.

FIG. 6 is a perspective view of the stalk which engages and disengages the canister in the inhaler of FIG. 5.

FIG. 7 is a side elevational view, partly in cross-section, of yet another embodiment of the inhaler, showing an alternative arrangement for achieving successive inhalations of a canister responsive to a patient's breath.

Detailed Description of the Drawing

FIGS. 1-3 depict a first embodiment of an inhaler 10 in accordance with the principles of the invention. The inhaler 10 includes a housing 12 which has a receptacle portion 14 connected to a mouthpiece 16. The receptacle portion 14 is in the

- 3 form of a sleeve shaped to receive a constant pressure canister 18 containing a medication. The canister 18 forms no part of the present invention. The inhaler apparatus of the present invention is made with any standard pressurized canister having an aerosol dispensing valve with a hollow discharge area which may be depressed manually with respect to the canister body area as inspiratory position is
- 10 which discharge of medication is presented, or an expiratory position in which a measured volume of the canister contents is discharged through the hollow discharge area.

The mouthpiece 16 includes an open end 22 spaced from the receptacle portion 14, and a closed end 24 defined by an end wall 26 which is connected to the

- 13 receptacle portion 14. The end wall 26 is generally a generally rounded or hemispherical in shape, with an open end 28 and a closed portion of the end wall 26 defining the open end 22.

With reference to FIG. 3, the housing 12 further includes an annular

- 16 and nozzle assembly 29 supported by the end wall 26. The annular and nozzle assembly 29 includes a bore 31 which is adapted to receive the hollow discharge area (not shown in FIGS. 1-3) of the canister 18, and a nozzle discharge outlet 30 in fluid

communication with the hole 22. The nozzle discharge outlet 30 is subsequently located in the open end 24 and oriented so that no aerosol particles generally along the nozzle longitudinal axis 22 of the nozzle. The outlet 30 preferably has an inward deflection at the exit of less than about 0.025 inch, and more preferably between about 0.020 inch and about 0.025 inch.

Thus, upon the nozzle 10 being depressed in the downward direction in FIG. 1, a normal vector of acceleration will be discharged from the hole 22 and into the outlet 30 to form a generally radial plume of accelerated particles within the nozzle 10, directed generally toward the open end 20 thereof. The outlet 10 includes features which promote dispersion and mixing of the accelerated particles via air which is emitted to reduce impingement and decrease the velocity of the liquid projectiles discharged from the nozzle 10. More specifically, the outlet 10 includes an air slot 34 disposed within the nozzle 10. The air slot 34 has an outlet 36 which is spaced downstream of and in opposing relationship with the nozzle 30 which is spaced downstream of and in opposition relationship with the nozzle outlet 22, and an inlet 38 which is in fluid communication with another air outlet 40 within the nozzle 10. In the configuration shown in FIGS. 1-3, the air slot 34 is a baffle valve which has a generally radial portion 40 which is generally aligned along the nozzle's longitudinal axis 22, and a generally radial portion 42 which is oriented in an angle with 44 of the nozzle 10. When a user exerts no impingement effort on the open end 20 of the nozzle 10, air is drawn from within the nozzle 10 into the air inlet 38, exiting the air slot outlet 34 in a direction toward the nozzle discharge

outlet 30. The portion 40 of air slot 34 is located and oriented within the nozzle 10 so that air flowing out from the outlet 34 will impinge on a plume of aerosol exiting the nozzle outlet 30. Once this air flows down the slot 34 has been established, the resulting valve of the nozzle 10 is actuated to discharge a plume of accelerated

particles from the nozzle 10. The impingement of air from air slot 34 on the plume causes the plume to slow down and be dispersed so as to occupy a larger portion of the cross section of the nozzle 10. The result is enhanced mixing of the aerosol with air, which promotes more complete vaporization of liquid projectiles by the time the aerosol leaves both the open end 20 of the nozzle 10, and a reduction in velocity of the plume exiting the open end 20 so as to approach the velocity of the impingement baffle. Accordingly, a greater fraction of the aerosol does of acceleration dispersed from the nozzle 10 exits the open end 20 in the form of respirable dry particles of the optimum size of about one to three microns having a relatively low velocity and substantially matching the impingement baffle velocity, as opposed to relatively large liquid droplets having a relatively high velocity, dispersion and mixing of particles within the nozzle and throat are thereby enhanced.

The air slot 34 and nozzle 10 can be integrally formed of one piece, with the inserted passage of the air slot 34 connecting through the nozzle 10 to another fluid communication with air within the nozzle 10. Alternatively, the air slot 34 can be formed of a small tube bent into the appropriate configuration and attached to the nozzle 10 at the inlet end 38.

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ALTHOUGH THE COUNTERCURRENT CHANNEL IN FIGS. 1-3 AND 7 SHOWS THE AIR SLOT 34 having an angle of 90 degrees w.r.t. the nozzle 40, it would be possible to have an angle of 45 degrees or less, or even greater, as long as the angle is not so great as to interfere with the advantages of the invention. For example, an angle of 45 degrees may be arranged in an acute angle (i.e., between about 10 degrees and 120 degrees), 120 degrees being defined as exactly opposite to the direction of a plume exiting the nozzle 10 or the exit 41 of the nozzle outlet 30, with the portion 40 of air slot 34 being oriented in direct or air jet at the nozzle 10. Additionally, the portion 42 which attaches to the nozzle wall can be varied, but can be oriented in an acute or obtuse angle to the nozzle wall 44.

The inventive feature includes features which reduce the likelihood of liquid droplets or dry particles impacting and permanently sticking to the lower walls 36 and 44 of the nozzle 10. More specifically, the outlet 10 includes a plurality of auxiliary air slots 46 through the end wall 34 and circumferentially spaced downstream of a lower air slot 34 from the nozzle outlet 30. A first circumferential ring of auxiliary air slots 46 are located adjacent to junction 42 between the end wall 34 and the lower wall 44 of the nozzle 10. A second circumferential ring of auxiliary air slots 47 are located centrally between the junction 42 and the nozzle outlet 30. An impingement effort caused on the open end 20 of the nozzle 10 causes air to flow into the auxiliary air slots 46 and 47 as indicated by arrows 25, and impinges therewith along the lower wall 44 of the nozzle 10 and

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against lower wall 34, as indicated by arrows 23. This auxiliary air flow forms a buffer or boundary layer air flow along the lower wall 44 and end wall 34 which tends to reduce the impact and permanent sticking of particles on lower wall 44 and end wall 34.

To the further advantage of this art, the outlet 10 also includes a plurality of venturi passages or vents 26 (also seen in FIG. 2) arranged on the lower wall 44 of the nozzle 10 and extending laterally therefrom. The vents 26 are located downstream of the auxiliary air slots 46, with each vent 26 substantially being located approximately in axial alignment with one of the auxiliary air slots 46. The vents 26 are oriented at an angle to the axial direction defined by longitudinal axis 22, so that venturi and vent 26 are oriented to be flowing over them. Thus, the boundary layer air flow caused by auxiliary air slots 46 causes the vents 26, which impinge venturi and vent 26 in the boundary layer air flow. This venturi and vent 26 further reduces the likelihood of aerosol droplets or particles impacting and permanently sticking to the lower wall 44.

As shown in FIGS. 1 and 3, the outlet 10 includes a separate nozzleplate 28 which connects to the open end 20 of the nozzle 10. The nozzleplate 28 has a central circular aperture 30 adapted to be inserted into the mouth of a user of the outlet 10. After completely extracting the user leaves the nozzle 28 but the mouth with the lips closed around the nozzle 28, and then begins to inhale, which establishes air flow from the air slot 34 and through the auxiliary air slots 46. Once

down air flow or established and with continuing to blade. On one side of the section 13 to discharge a second velocity of medication and propellant mixture from the nozzle discharge outlet 30. The user continues to turn to 13 for longer to mix capacity, and thus typically holds the handle for a period of time to allow the accumulated materials in nozzle within the storage of the bags.

- As shown in FIGS. 1-3, the housing 12 is formed in four sections containing the components 30 which subsequently fit together. However, for ease of assembly, the housing 12 may alternatively be formed in fewer than four sections. For example, the housing 12 may be formed in two sections, a first section 10 including the nozzle portion 14, and wall 24, and the nozzle 15 up to and including the valve 24, and a second section including the portion of nozzle 15 leading the air into 24 and the component 30. Alternatively, the housing 12 may be formed in two sections split on a longitudinal plane through the nozzle, the two sections being generally either images of each other which are joined together along the plane of symmetry. Nevertheless, for illustrative purposes, an embodiment having four sections is shown and described.

- A first section 10 includes the nozzle portion 14, the end wall 24 and actuator and nozzle assembly 26, and a generally cylindrical portion 42 which forms a part of the nozzle 15 and is connected to the end wall 24 at the junction 48. The first section 10 advantageously is longitudinally formed of two pieces, although it may alternatively be formed in multiple pieces which are subsequently joined together.

A second section 44 includes a second generally cylindrical portion 46 which inner and outer diameters are equal to those of the first generally cylindrical portion 42, and a reduced-diameter portion 48 which is subsequently received within the longitudinal space end of first cylindrical portion 42. The portion 46 has an inner wall 50 which is generally coaxial, converging slightly in the axial direction toward the component 30. The valve 24 is mounted on the inner wall 50. Second section 44 preferably is longitudinally formed of one piece, although it may alternatively be formed in multiple pieces which are subsequently joined.

- A third section 72 of the housing 12 includes a third generally cylindrical portion 74 where inner and outer diameters are equal to those of the second generally cylindrical portion 46, and a reduced-diameter cylindrical portion 76 which is subsequently received within the axial downstream end of second generally cylindrical portion 46. The outer diameter of portion 76 is approximately equal to the inner diameter of portion 46 so as to provide a tight fit between these parts. The inner surface 78 of portion 76 has a diameter which is approximately equal to the smaller diameter of the nozzle inner wall 50 so that the junction between sections 70 and 72 does not present any substantial step in the flowpath defined by the nozzle 15. The air valve 34 is mounted on the inner surface of the third section 72 at the junction between the inner surface 78 and the inner surface 50 of third cylindrical portion 74. A hole 82 through the portion 74 connects with the internal passage of air valve 24 to provide fluid communication between the tube 18 of air valve 24 and

section 44 outside the nozzle 15. Third section 72 may be longitudinally formed of one piece, or formed in multiple pieces and subsequently joined.

- The fourth section of the housing 12 is the component 30, which has a generally cylindrical portion 34 which is subsequently received within the open downstream end of the third generally cylindrical portion 74 (which also defines the open end 32 of the nozzle 15). The portion 34 is attached to an annular flange 36, which is turn is attached to the reduced-diameter portion 28 which is inserted into a user's mouth. The outer diameter of portion 34 is approximately equal to the diameter of inner section 46 so as to provide a tight fit therebetween.

- The housing 12 advantageously is formed of a plastic such as polypropylene, polyurethane, polycarbonate, ABS, polychloroform, or polycarbonate. The housing 12 may be manufactured by any suitable technique such as injection molding or blow molding.

- FIG. 3A shows an alternative embodiment of an actuator and nozzle assembly 26 for the handle 10, in cross-sectional view on the horizontal plane indicated in FIG. 3. The actuator and nozzle assembly 26a includes two spaced-apart open discharge outlets 30a which are both directly connected to the base 26a and which converge toward each other in the direction of the component 30. Thus, depressing the handle 12 as to discharge a second velocity of medication into the base 26a causes two aerosol plumes to exit from the pair of outlets 30a. The plumes converge and impinge on each other upstream of the air valve outlet 24.

causing the aerosol or spray mix, thereby aiding mixing of the aerosol with air. Additionally, impingement of the two plumes aids in creating greater droplets, which enhances respiration of propellants. It will be appreciated that for convenience of illustration, the base 26a is shown as being elongated in the horizontal direction and section 30a are shown as being spaced apart in the horizontal plane. Alternatively, however, the base 26a may simply be extended in the vertical direction and the sections 30a vertically spaced apart and angled toward each other so as to achieve the desired convergence of the two plumes.

- FIG. 4 depicts an alternative embodiment of a handle 12b in which the elongated air valve 34 of handle 10 has been replaced by a shorter air valve in the form of a tube 42b which is supported in the handle 12b by a pair of bellows 48a. In FIG. 4, ports identified by reference numerals having the letter "A" indicate ports analogous to those having the same reference numerals within the text in FIG. 3, while ports identified with identical reference numerals in FIGS. 3 and 4 denote identical ports. Thus, the tube 42b is analogous to the actuator 42 of the air valve 24, and the cylinder 48a are analogous to the actuator 48 of air valve 24. The tube 42b includes a central cavity 43 of a low diameter, and an outer passage 46a of a much wider diameter. The outer passage 46a is generally coaxial with the handle 12b and oriented so that air flowing downward direction is directed toward the nozzle outlet 30. The internal passage of cylinder 48a are connected to provide air by a pair of tubes 42b through the cylindrical portion 74a. In the

construction of the inhaler 12 shown in FIG. 4, there is no outlet of the breathing tube 10 at the second section 34 of FIG. 3. Thus, the nozzle 34 here is disclosed from the inhaler 10. However, the nozzle or tube 40 are still present in the inhaler 12 to provide a breathing tube air flow along the front wall of the nozzle 10.

FIGS. 3 and 4 illustrate yet another construction of an inhaler 12 in accordance with the principles of the present invention. FIG. 3 schematically depicts a breathing canister assembly analogous to FIG. 1, showing an inhaler 12 in which the aerosol plume is directed away from the user so that the aerosol canister 10 is directed laterally. FIG. 4 schematically depicts a vertical canister section of the inhaler 12. Again, 120 puffs are issued by the inhaler canister, with discharge ports are shown by the letter "B" suffix. The inhaler 12 includes a housing 120 defining a nozzle 120 which has a flow channel 120 defined by an end wall 120 and a second space 120 defined by a nozzleplate portion 120 aligned to be directed toward a user's nose. The nozzle 120 has a flow larger internal cross section than the nozzle 10 of FIG. 1, according to a second smaller internal cross section area of the nozzleplate portion 120. The housing further includes a nozzleplate portion 120 which terminates just the nozzle 120 at a location between the end wall 120 and the nozzleplate portion 120. The nozzleplate portion 120 receives a standard pressurized canister (not shown). The housing 120 further includes an actuator and switch assembly 20 arranged at the bottom end of nozzleplate portion 120.

such that the bottom ends area of the canister may be inserted into a base 20 of the actuator and switch assembly 20. The body of the actuator and switch assembly 20 have already been described in connection with FIG. 3. The aerosol discharge outlet 30 is oriented so as to direct an aerosol plume toward the end wall 120.

5 The inhaler 120 includes an inhaler canister 12 which is directly disposed with the nozzle 120. The inhaler canister 12 has an open end 120 spaced from and adjacent the end wall 120, and a closed end 120 recessed from the end wall 120 and defined by an end wall 340 which separates the actuator and switch assembly 20. The inhaler further includes an air tube 340 attached to the end wall 120 and centrally disposed within the nozzle 120. The air tube 340 further provides way for the user nozzle 120 toward the aerosol discharge outlet 30. The base 20 of the actuator 20 is connected to ambient air through the nozzle 120 by a tube 30 through end wall 30. The nozzle 30 of air tube 340 is in opposing relation to the nozzle 30. Airflow originates from the nozzle 30 passes into the interior of base canister 20 and proceeds toward the end wall 30 of base canister 20. Inhalation of air through the nozzleplate 30 causes air to enter through tube 30 into air tube 340 and out the nozzle 340 toward the plume. The plume and air jet from the nozzle 340, causing the plume to slow down and spread out within base canister 20. Continued inhalation by the user causes the displaced aerosol to exit through the open end 120 of base canister 20, and then reverse directions to flow through the space between the base canister 20 and the user nozzle 120, and then through the nozzleplate 30.

Thus, the aerosol reaches a portion of the length of nozzle 120 earlier, thereby increasing residence time of the aerosol within the device before exiting the nozzleplate 30. This leads to more complete expansion of liquid droplets. Furthermore, the user aerosol knows that the velocity of the aerosol exiting the nozzleplate will be substantially equal to the velocity of the user's inspired breath, reducing the potential of impaction in the mouth and throat.

FIG. 7 depicts yet another construction of the inhaler providing automatic extension of the canister to discharge a dose of medication in response to an predetermined width, the user's respiratory effort. An inhaler 120 includes a housing 120 having a nozzle 120 which an aerosol plume is caused to originate by the user. The nozzle 120 is shown to include the air tube 34 and the nozzle air tube 40. It may also include the nozzle 30 of inhaler 10. Alternatively, the nozzle 120 may be a simple nozzle tube with an open end for the exit of aerosol particles. Thus, with the exception that the nozzle 120 may be aligned to provide field encompasses with a chamber 120 is housing 120 is disclosed below, the details of the nozzle 120 are not important in an understanding of the breath synchronization feature of the inhaler.

The housing 120 further includes a nozzleplate portion 140 which is connected to the nozzle 120. The nozzleplate portion 140 comprises a generally cylindrical shape having a longitudinal axis 120 which is angled at an oblique angle to the longitudinal axis of the nozzle 120. A chamber 120 resides within the nozzleplate

portion 140 with its longitudinal axis aligned with the longitudinal axis of the nozzleplate portion 140. Disposed between the nozzleplate portion 140 and the nozzle 120 is an inner sleeve 120. The inner sleeve 120 has an open top end 120 through which the canister 12 may be inserted, and an open bottom end 120 which is connected such that the canister 12 passes through it in which connection provides the bottom area 120 of the canister to be inserted into the base 20 of actuator and switch assembly 20. More specifically, the base 120 adjacent bottom end 120 has inward extending legs 120 which seat on cap portion 120 of the canister. The canister 12 is stable within base 120 during the direction defined by the longitudinal axis 120 of nozzleplate portion 140 so as to prevent the canister 12 from disengaged toward the actuator and switch assembly 20 in order to estimate the canister's operating valve.

The inner sleeve 120 is also stable within the nozzleplate portion 140 along the direction of axis 120 for the purpose of placing the canister 12 in a correct position ready to be activated. The nozzleplate portion 140 has two longitudinal axis 120 circumferentially spaced apart 180 degrees, one of which includes a pair of diametrically opposite legs or ears 120 extending outwardly from the outer surface of inner sleeve 120. Alternatively, the nozzleplate portion 140 may have only one axis 120 spaced 180 degrees apart and including the legs 120. Thus, as the base 120 moves laterally longitudinally within nozzleplate portion 140, the legs 120 slide longitudinally within the respective slots 120.

The holder includes a generally cylindrical case cap 114 which fits over the cap of receptacle portion 14a. The case cap 114 has an annular flange 116 at its lower end which extends outward beyond the outer surface of the housing so as to facilitate gripping of the case cap 114 by the user's hand. The lower surface 118 of case cap 114 has a pair of circumferentially extending recesses or case seats 120 formed thereon approximately 120 degrees apart which extend longitudinally upward in the open top end 122 of case cap 114. Each case seat 120 receives a generally U-shaped sleeve 124 in facing relationship with one of the legs 112 protruding downward from the lower surface 120 through slot 116. Thus, starting with the case cap 114 in a position in which each leg 112 is in contact with the lowermost portion of the respective case track 120 (i.e., the portion of case track 120 which is furthest from the top end 122 of case cap 114), rotation of the case cap 114 through the arc defined by the case seats 120 causes the legs 124 to ride along the U-shaped surfaces 120 and thereby gradually advances the lower sleeve 124 in the longitudinal direction toward the top end 122.

This gradual advancement of the lower sleeve 124 drives the cylinder 12 through rotation of the hub 122. Rotating the upper portion of the cylinder 12 is a compression spring 126. The spring 126 is attached to the lower surface of a removable cap 128 which surrounds the top end 122 of the receptacle portion 14a and the top end 122 of the case cap 114 to completely enclose the cylinder 12 in the housing. When the cap 128 is thus breached, the spring 126 bears against the end

of the cylinder 12. During the cylinder movement toward the bottom end nozzle assembly 24, this bearing or impacts the downward movement of the cylinder 12. The spring 126 would move the cylinder downward until the discharge area 19 were fully depressed from the cylinder so as to cause discharge of a measured volume of the cylinder contents. However, the holder 12 includes a mechanism which prevents the cylinder to prevent this downward movement, with the mechanism being responsive to an impingement effort of a sleeve carried in the open end of the nozzle 14a so as to disengage from the cylinder during the user's intention to allow the spring 126 to move the cylinder from its discharge position.

To this end, the holder 12 includes a plunger assembly 132 which is rotatable relative to the cylinder 12 along an axis 124 generally parallel to the longitudinal axis 120. The plunger assembly 132 includes a circular disc 134 having a slot 136 extending centrally therethrough provided with ends 138 and presenting outward from both sides of the disc 134. A first portion 140 of the slot 136 presenting from the side of the disc 134 remote from the cylinder receives a sleeve 142 in a slot 144 of the housing, the sleeve 142 guiding the movement of the plunger assembly 132 along axis 124. A second portion 146 of slot 136 presenting from the side of disc 134 holds the cylinder catch through a opening 148 in receptacle portion 14a, functioning as an enlarged head end 132. A compression spring 126 is capture between the head end 132 and the wall of the receptacle portion 14a, holding the plunger assembly 132 toward the cylinder 12.

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A latched trigger 154 is attached to the head end 132. The trigger 154 has two spaced-apart parallel prongs 154a, b which extend along the direction of axis 124 in approximately the longitudinal axis 120 of the receptacle portion 14a. The prong 154a is spaced apart by a distance > width of a slightly smaller than the diameter of the cylinder neck 123 from which the discharge area 19 protrudes, so shown schematically in FIG. 8. Thus, when the plunger assembly 132 is fully extended toward the cylinder 12, the cylinder neck 123 receives lower edge portions 140 of the prong 154a, or held firmly by the slanted surfaces in FIG. 8. However, when the plunger assembly 132 is withdrawn along axis 124 away from the cylinder 12, the cylinder neck 123 does not engage 154a as the consequence of the cylinder 12 moved the sleeve 142 to a proximal end. The prong 154b includes portion 157 which sharply curves away from the cylinder neck 123 in the direction along axis 124 toward the cylinder. The portion 157 carries the elements of force required for disengagement of the trigger 154 from the cylinder neck 123.

Movement of the plunger assembly 132 in the direction away from the cylinder is responsive to air pressure within a variable-volume chamber 162 within the housing. The chamber 162 is defined by the side 126, the housing wall 144, and a flexible diaphragm 164 which separates the side 126 to the wall 144 in a substantially air-tight manner. The diaphragm 164 includes a diaphragm portion 166 which lies upon the side of side 126 facing the cylinder 12, and a skirt 168 which departs from the lower edge of the diaphragm 164 and extends to the housing

wall 144. Further advancement, the housing wall 144 receives a removable cover 170 of the housing, and an edge of the skirt 168 is attached to the housing by being snap-fitted between the cover 170 and the remainder of the housing. The diaphragm portion 166 of diaphragm 164 includes a central hole through which the side 126 extends and which slightly increases the skirt 168 to provide a substantially air-tight seal therewith.

The removable cover 170 includes a passage 172 facing the side 126 which aligns with a passage 174 defined in a ribbed 176 of the housing. The passage 174 extends toward the open end 122 of receptacle 14a. The ribbed 176 is defined in a lower free surface, a thin generally cylindrical section 178 which includes the ribbed 176 and is connected to the end wall 124 through which the flexible section 166 extends, and a raised generally cylindrical section 180 which includes the air valve 14 and which connects to the first section 176. The passage 174 terminates at the end of first section 176 which connects to second section 178. A passage 178 through a ribbed 180 of the second section 178 is directly connected with and forms an extension of passage 174. The passage 178 extends from the forward passage 172 of the air valve 14. A ribbed 182 is inserted into the air valve passage 172. The ribbed 182 includes a ribbed portion or skirt 184. Air passage 182 extends through the ribbed 182 in the vicinity of the diaphragm 164. The ribbed 182 is disposed in passage 180 such that there air passage 182 aligns with the passage 178. Thus, fluid communication is provided between the second section 178

and the venturi-vortex chamber 122 by air pressure 123, pressure 124 is across section 204, pressure 125 is after section 205, and section 222 is after 170.

At 440 Cycles per second the velocity in the venturi chamber 122 is approximately 100 ft/sec and 20% of ambient air, air is drawn from outside the chamber 122 through air tube 34 has the primary air pressure of the chamber 122. This air is then drawn through the venturi 124, and consequently a below-atmospheric pressure exists in the venturi 124. This below-atmospheric air pressure is communicated to the chamber 122, with the result that the walls of the chamber 122 are subjected to a force proportional to the pressure difference between atmospheric pressure outside the chamber 122 and the below-atmospheric pressure inside the chamber 122.

Consequently, as within the chamber 122 begins to increase the chamber 122 through section 172, through pressure 174 and 175, through pressure 123, and has the venturi section 124, and thence through the air tube 34 has the primary air pressure of the chamber 122.

At the new conditions of intake through the venturi 124, expansion of air from the chamber 122 causes the velocity in chamber 122 to increase, with the result that the air 126 and air shock 223 begin to move around the vent 124 against the force of the spring 125. Accordingly, the trigger 124 begins to move so as to discharge the pressure 123 from the chamber vent 124. When the decrease in volume is sufficient to move the trigger 124 far enough to easily discharge the pressure 123 from the vent 124, movement of the chamber 122 around the venturi 124 is no longer

impeded, and the force of spring 125 causes the chamber movement so as to cause extension of the chamber's venturi valve. A certain rate of venturi extension is thereby discharged from vent 124 into the venturi 122 for initiation by the user.

After the initiator 120 has been activated to dispense a dose of medication, it must be reseated so that it is ready to be discharged again. To this end, the user grasps the ring 114 and rotates it with respect to the housing 121 through the slot defined by the slot notch 221. This causes the lower sleeve 120 and section 15 to be rotated upward against the force of spring 125. When the chamber 122 is rotated sufficiently clockwise so that the trigger 124 is clear the chamber vent 124, the trigger 124 begins to move toward the chamber 122 so that the trigger 124 once again is in a fully extended position to engage the chamber vent 124. The user then rotates the outer ring 114 back to its starting position to lower the chamber 122, whereupon the chamber vent 124 once again opens the pressure 123 of the trigger 124. The initiator 120 is then ready to be used again.

It will be appreciated that the breath-expansion devices described above provide an initiator in which discharge of medication is automatically responsive to the user's respiratory effort, so that the user does not have to manually commence manual depression of a chamber with the initiator. Furthermore, discharge of medication does not occur immediately upon the user beginning to inhale on the open end of the device, but rather is suspended until the volume of chamber 122

has decreased enough to close sections. It will also be appreciated that the degree of flow delay between initiation of a breath and sections is dependent on a number of factors, the primary factor being the cross-sectional area of the chamber 122 and the spring constant of the spring 125, since a discharge of medication requires a certain minimum level of the chamber 122 to cause the discharge area 13 to be fully depressed, and the travel is proportional to the pressure difference across the chamber 122 as cross-sectional area divided by the spring constant. Accordingly, the initiator 120 may be designed with appropriate selection of these factors so as to achieve closure of the chamber 122 near the peak of a user's inhalation.

Moreover, the initiator 120 provides breath-expansive amounts of the chamber 122 which automatically adjust to the user's rate of inhalation to discharge the medication near the peak of the inhalation, i.e., near the point at which 50 percent of the volume which the user will ultimately inhale with a full inhalation has been inspired. For instance, if a user with normal lung function inhales quickly through the open end 204, air will be removed from the chamber 122 more rapidly so as to achieve sections in a relatively short time. Conversely, if a user with impaired lung function inhales slowly through the open end 204, air will be removed more slowly from chamber 122 so as to achieve sections in a relatively longer time.

The initiator 120 further includes an adjustment screw 220 which extends through the housing 121 has the pressure 174 to limit a restriction within pressure 174. By rotating the screw 220 one direction, the screw 220 causes further

low pressure 174 to increase the restriction, and by rotating the screw 220 in opposite direction, it restricts to decrease the restriction. Thus, the setting of sections of the chamber 122 in relation to a particular patient's inhalation may be varied by adjusting the screw 220. Varying the screw position results in a variation in pressure

difference across the walls of the venturi-vortex chamber 122 as a given flow rate over the open end 204 of chamber 122. Thus, for a given flow rate over the open end 204 of chamber 122, setting the screw 220 to increase the restriction of pressure 174 will increase the flow period required to overcome the chamber 122 sufficiently to cause sections, whereas setting the screw 220 to decrease the restriction will decrease such flow period.

FIG. 9 depicts a substantially improved initiator embodiment of an initiator having features for automatic breath sections of discharge. In this embodiment, the initiator trigger 124 is dislodged and the displacement plunger assembly 121 is replaced by a mechanically compressible bellows 320 which is disposed between a front wall 321 and rear wall 324 of the initiator 120. The initiator 120 has a base 322 as the exterior which holds the initiator in a conventional position, the initiator being supported by air pressure from a position permitting the initiator to move from a discharge position.

The bellows 320 is substantially made of resilient material and has a flat end 323 or the end adjacent the initiator vent 124, the flat end 323 being integrally formed with the initiator 120 in a conventional state with 320. The bellows 320 has a

bottom and well 220 as the air enters the bottom wall 221, the air and well 220 being largely defined with the air and well 221. The bottom and well 220 is placed by a side or inside 220 which constitutes an air passage into the interior of the bottom 221. The nozzle 219 subsequently is a nozzle and also similar to a hypodermic needle and is largely defined as one end to the air and well 220 by meeting at other nozzle sections. The air and 222 of the nozzle 220 extends to another air nozzle 219 to the air and well 221 of a nozzle 216. The nozzle 216 is disposed within a side 216 which extends down as side 220 which draws air from outside the bottom housing, as an air and well 222 which is arranged within the nozzle 216 that thereof appears the nozzle discharge article 20. The side 216 and nozzle 216 may also be formed of suitable metal.

A supportable plenum 224 is attached to the bottom and well 220 of the bottom 221. The supportable plenum 224 receives the nozzle such 125 throughout the range of nozzle discharge by the nozzle in moving from a rest or ready position to a discharge position. The bottom 221, via the supportable plenum 224, carries a spring force on the nozzle such 125. The force of the bottom 220 acts in a direction tending to move the nozzle such 125 away from the nozzle 216. Additionally, as is well known, the nozzle 125 undergoes an internal spring (not shown) which acts between the nozzle body and the bottom nozzle cap 17 in a direction tending to move the nozzle 125 away from the nozzle 216. The spring means of the bottom 220 is selected such that the sum of the spring force

exerted by the bottom 220 and the force caused by the bottom spring is slightly greater than the force caused by the spring 125 (FIG. 7) which exerts a force on the end of the nozzle 125 in the direction as tend to move the nozzle 125 toward the bottom 216 less its discharge position. Thus, at rest, with atmospheric pressure acting both inside and outside the bottom 220, the bottom 220 and bottom spring overcome the force of the spring 125 and thereby keep the nozzle 125 in a ready position preventing discharge of medication therefrom.

However, when a user breathes through the nozzle (not shown) of the inhaler, air is drawn through the side 216, as previously described in connection with the bottom 216, which creates a low pressure within the air and well 216 of nozzle 216. This low pressure is communicated via the nozzle side 212 and nozzle 220 to the interior of the bottom 220. As a result, the pressure within the bottom 220 is less than the atmospheric pressure which overcomes the nozzle of the bottom 220, and therefore there is an air pressure force applied on the bottom and well 220 in the direction toward the bottom wall 221. The sum of this air pressure force and the force of the spring 125 exceeds the spring force exerted by the bottom 220 and so causes internal spring, causing the bottom and well 220 of bottom 220 to be compressed toward the bottom wall 221. By virtue of the forces caused on the nozzle 125 by the spring 125, the nozzle 125 is moved into its discharge position 20, evacuation of air from the bottom 220, the nozzle 125 is moved into its discharge position. Once the user completes his inhalation and air flow through the nozzle 216

ceases, air pressure is again equalized inside and outside the bottom 220, and the bottom 220 returns to its starting position, the force of the bottom 220 and bottom spring tending the nozzle 125 back toward spares the force of the spring 125 from the ready position. Thus, with the breath-activated system depicted in FIG. 7, there is no need for a separate switching device.

The bottom 220 preferably has a spring constant of about 1 pound per inch to about 17 pounds per inch, and a cross-sectional area of about 0.1 to about 0.75 square inch. Thus, a pressure differential of about one pound per square inch across the bottom 220 is sufficient to compress the bottom 220 by an amount of about 0.025 inch to about 0.020 inch. With a standard nozzle 125, only about 0.005 inch of relative movement is required between the discharge nozzle 125 and the nozzle body in order to cause discharge. Accordingly, the nozzle 216 must be close to cause a gap pressure which the air and well 221 of about one pound per square inch.

With the present invention has been illustrated by a description of various embodiments and while these embodiments have been described in considerable detail, it is not the intention of the applicants to restrict or in any way limit the scope of the appended claims to such detail or to such embodiment. For example, while the inhaler which is claimed and described herein for aerosol inhaler is communicated with ambient air via a passage through the nozzle well, the aerosol inhaler may alternatively draw air through one of the auxiliary air holes 40 in the side wall 34, or

through any arrangement having the same like results for primary air passage defined by the inhaler nozzle. Additionally, the nozzle used between 200 of FIG. 8 may advantageously be used in the inhaler configuration depicted in FIG. 7, with the bottom 220 replacing the plenum assembly 122 and the bottom and well 220 of the bottom 220 being attached to the bottom plenum 124, and the spring 125 being substituted by virtue of the stiffness of the bottom 220. The inhaler is to broader aspects is therefore not limited to the specific details, representations, apparatus and methods, and illustrative examples above and described. Accordingly, departure may be made from such details without departing from the spirit or scope of applicants' general inventive concept.

With reference to the use of the words "aerosol" or "aerosolized" or "aerosolizing" in the foregoing description neither in the following claims, nor in the claims against which reference is made, are the words to be interpreted literally, either individually, and/or each of these words is to be an ingredient to encompassing the foregoing description under the following claims.

The claims defining the invention are as follows:

1. An aerosol flow control apparatus providing automatic discharge of medication responsive to an inspiratory effort of a user, the apparatus comprising:
a pressurized canister containing a canister body and a bottom discharge area which is movable with respect to the canister body between an inspiratory position in which discharge of medication is prevented and an expiratory position in which medication is discharged through the discharge area;
a housing adapted to support the canister and permit movement thereof between a first position in which the discharge area is in the inspiratory position and a second position in which the discharge area is in the expiratory position, the housing further defining a primary air passage including an outlet through which a user can inhale and also defining a secondary air passage connecting the primary air passage and canister or canister discharge area; and
a variable-volume device supported within the housing and including a wall which is movable with respect to the housing, the variable-volume device defining a variable-volume chamber having a fluid communication with the variable device;
a canister carrier affixed to the variable wall of the variable-volume device, the canister carrier being movable with the variable wall from a first position in which the canister is in the first position and another movement between the canister body and discharge area is prevented, to a discharge position in which the canister is free to move into the second position;
a main body portion which urges the canister into the second position upon movement of the canister carrier into its discharge position; and
the variable-volume chamber being in fluid communication with the primary air passage, whereby inhalation of a user through the outlet causes air to be drawn through the variable device causing a low pressure in the throat which is communicated to the variable-volume chamber, the low pressure causing air to be evacuated from the chamber and thereby cause the variable wall to move the canister carrier into the discharge position.
2. The aerosol flow control apparatus of claim 1, wherein the variable device is connected to the chamber by a third air passage within the housing, and further comprising an adjustment device which may be selectively positioned to selectively vary the flow rate

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through the third air passage at a given flow rate through the primary air passage, thereby varying the timing of medication discharge in relation to the breathing cycle of a user.

3. The aerosol flow control apparatus of claim 1, wherein the variable-volume device comprises a plenum which is rigidly connected to a wall of the housing by a flexible diaphragm, and the canister carrier includes a member which is attached to the plenum and which in the first position extends into the path travelled by the canister between the first and second positions so as to prevent the canister from moving from the second position, evacuation of air from within the chamber of the variable-volume device causing the plenum to move toward the housing wall and thereby release the member from the discharge position permitting the canister to move into the second position.
4. The aerosol flow control apparatus of claim 1, wherein the housing comprises a main body portion which encloses the canister, and an end cap which covers the end of the canister opposite from the end with the discharge area and which engages the main body portion to prevent inadvertent removal therefrom, the main body portion comprising a compression spring between an inner surface of the end cap and the canister such that the spring bears against the canister when the end cap is engaged with the main body portion.
5. The aerosol flow control apparatus of claim 1, wherein the main body portion includes a generally cylindrical receptacle having a longitudinal side and defining a generally cylindrical recess in which the canister resides, and further comprising a sealing device including:
an inner sleeve which surrounds the canister within the receptacle, the inner sleeve and canister being slidable together as a unit within the receptacle along the longitudinal side, the inner sleeve further including at least one pin extending outwardly from an outer surface thereof through a slot in the receptacle; and
a sealing ring which surrounds the receptacle and has a surface which engages the at least one pin, the sealing ring being movable with respect to the receptacle so as to move the pin in the direction defined by the longitudinal side toward the end cap so as to draw the inner sleeve and canister apertured and thereby move the canister into a closed position which permits the canister to move in and out of the receptacle, thereby modifying the apparatus for inhalation in response to the inspiratory effort of a user.

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open a user breathing through the outlet, drawing air through a secondary air passage arranged within the housing, the secondary air passage extending from the primary air passage to admit air outside the primary air passage, the secondary air passage including a vented housing channel; and

a slot during the drawing step, providing fluid communication between the draw position of the vent of the secondary air passage and the variable-volume air chamber so as to communicate a below-atmospheric air pressure caused by the vent to the air chamber and thereby cause the chamber volume to decrease, whereby the canister automatically moves to prevent said movement of the canister into the second position to discharge medication when the predetermined decrease in chamber volume is reached.

6. The method of claim 7 wherein the draw position of the vent has a reduced cross-sectional flow area relative to the vent of the secondary air passage such that the air pressure in the draw position is lower than the air pressure in the vent of the secondary air passage when air is drawing therethrough.

DATED Oct 14, 1986

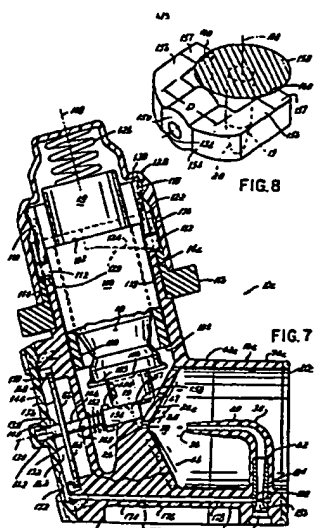
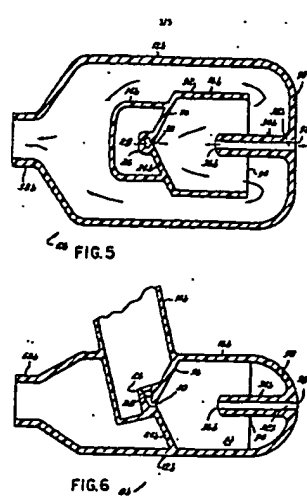
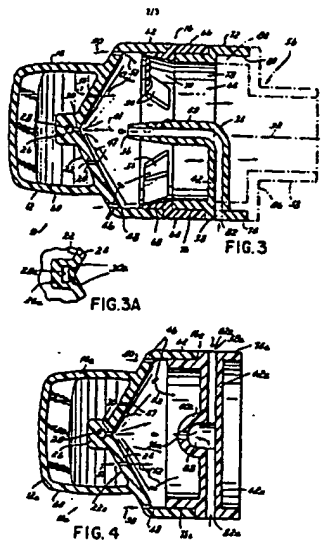
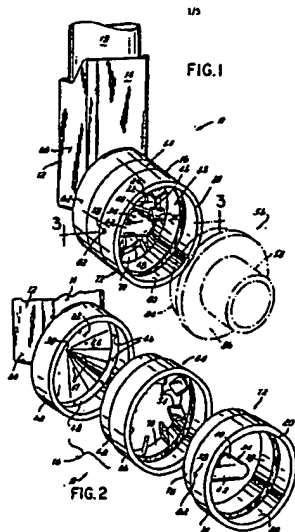
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7. A method of delivering a dose of medication using an aerosol delivery apparatus which houses a medication-containing canister having a canister body and a bottom outlet area which is movable with respect to the canister body between an inspiratory position in which discharge of medication is prevented and an expiratory position in which medication is discharged through the outlet area, with the canister being movable within the apparatus between a first position in which the outlet area is in the inspiratory position and a second position in which the outlet area is in the expiratory position, the apparatus including a housing defining a primary air passage having an outlet through which a user can inhale and a secondary air passage, the simultaneous change of medication from the canister with an inspiratory effort of a user through the outlet, the method comprising:
placing the canister in the first position;
preventing movement of the canister into the second position by a canister carrier which engages the canister to prevent said movement and which is movable in response to below-atmospheric air pressure within a variable-volume device arranged within the housing, the variable-volume device including an air chamber therein, the canister carrier being movable to permit the canister to move into the second position upon a predetermined decrease in volume of the air chamber;
engaging the canister toward the second position;

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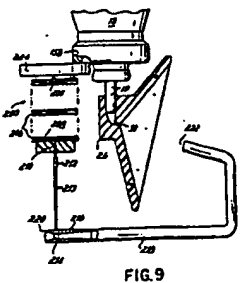


FIG.9